

Clinal variations in seedling traits and responses to water availability correspond to seed-source environmental gradients in a foundational dryland tree species

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• **Background and Aims** In dryland ecosystems, conifer species are threatened by more frequent and severe droughts, which can push species beyond their physiological limits. Adequate seedling establishment will be critical for future resilience to global change. We used a common garden glasshouse experiment to determine how seedling functional trait expression and plasticity varied among seed sources in response to a gradient of water availability, focusing on a foundational dryland tree species of the western USA, *Pinus monophylla*. We hypothesized that the expression of growth-related seedling traits would show patterns consistent with local adaptation, given clinal variation among seed source environments.

• **Methods** We collected *P. monophylla* seeds from 23 sites distributed across rangewide gradients of aridity and seasonal moisture availability. A total of 3320 seedlings were propagated with four watering treatments representing progressively decreasing water availability. Above- and below-ground growth-related traits of first-year seedlings were measured. Trait values and trait plasticity, here representing the degree of variation among watering treatments, were modelled as a function of watering treatment and environmental conditions at the seed source locations (i.e. water availability, precipitation seasonality).

• **Key Results** We found that, under all treatments, seedlings from more arid climates had larger above- and below-ground biomass compared to seedlings from sites experiencing lower growing-season water limitation, even after accounting for differences in seed size. Additionally, trait plasticity in response to watering treatments was greatest for seedlings from summer-wet sites that experience periodic monsoonal rain events.

• **Conclusions** Our results show that *P. monophylla* seedlings respond to drought through plasticity in multiple traits, but variation in trait responses suggests that different populations are likely to respond uniquely to changes in local climate. Such trait diversity will probably influence the potential for future seedling recruitment in woodlands that are projected to experience extensive drought-related tree mortality.

Key words: Common garden, forest and woodland recovery, drought effects, dryland woodlands, phenotypic plasticity, *Pinus monophylla*, pinyon pine, precipitation seasonality, intraspecific trait variation, seedling traits, tree regeneration.

INTRODUCTION

Seedling recruitment is critical for the long-term persistence of tree species (Lloret *et al.*, 2009), yet recruitment is often considered a bottleneck for plant populations due to high rates of mortality during the first months following emergence (Fenner, 1987; Lloret *et al.*, 2004). Having minimal carbon storage capacity and limited root systems, seedlings are more vulnerable to climate variability than mature plants (Lenoir *et al.*, 2009; Lloret *et al.*, 2009; Matías *et al.*, 2017). Interacting climate change factors have amplified both climate variability and the severity and duration of droughts (Niinemets, 2010), causing tree mortality worldwide (Allen *et al.*, 2010; Anderegg *et al.*, 2019). Forest and woodland ecosystem recovery depends on future stand replacement by seedling establishment (Redmond

et al., 2015; Andivia *et al.*, 2018). Thus, understanding early-stage phenotypic traits that enhance survival across environmental gradients can inform predictions of forest resilience to rapidly changing conditions (Ramírez-Valiente *et al.*, 2021).

Tree seedling growth-related traits, both above and below ground, influence drought tolerance by affecting seedlings' ability to acquire and store resources (Sánchez-Gómez *et al.*, 2010). Many studies on pine species have found that seedlings from drier climates invest more growth in their roots when water resources are limited (Richter *et al.*, 2012; Moser *et al.*, 2015; Kolb *et al.*, 2016), and grow fewer needles to reduce water loss (Sánchez-Gómez *et al.*, 2010). Increasing the investment in roots relative to shoots (i.e. higher root mass fraction; increased root biomass allocation) can enhance access to water

and nutrients (Markesteijn and Poorter, 2009; Poorter *et al.*, 2012; Freschet *et al.*, 2015). Likewise, high specific root length, or the ratio of root length to dry mass of roots, tends to increase uptake rates and is considered the below-ground analogue for specific leaf area (Ostonen *et al.*, 2007).

Dryland tree species inhabit environments that are water limited with high interannual precipitation variability and are characterized by infrequent, episodic precipitation events during the growing season (D’Odorico and Bhattachan, 2012). Plastic phenotypic trait variation, the ability of an individual to alter its phenotype in response to the environment (Albert *et al.*, 2011; Klingenberg, 2019), represents an important strategy for coping with fluctuating conditions (Ghalambor *et al.*, 2007; Matesanz *et al.*, 2010; Nicotra *et al.*, 2010; Lázaro-Nogal *et al.*, 2015) and is known to have a heritable genetic basis (Scheiner, 1993). An adaptive strategy for seedlings of some species in variable environments is to increase above-ground growth during periods of high water availability while reducing growth during periods of water stress (Richter *et al.*, 2012). Episodic precipitation events can thus have significant effects on plant development and survival (Gillespie and Loik, 2004) and can contribute to the evolution of plasticity in particular seedling traits (Pratt and Mooney, 2013).

For species that occupy a large geographical range, differences in the environmental conditions experienced by distinct populations may influence trait values and plasticity patterns and produce locally adapted phenotypes (Lázaro-Nogal *et al.*, 2015). In general, populations experiencing the greatest environmental heterogeneity are expected to be more plastic in their traits, which may allow them to better manage abiotic stress (Pratt and Mooney, 2013; Valladares *et al.*, 2014; Lázaro-Nogal *et al.*, 2015). Populations occurring in more stressful locations, such as trailing-edge populations, have also been shown to have more plasticity (Chevin and Lande, 2011; Lázaro-Nogal *et al.*, 2015). The ability to quickly respond to changing conditions can confer a fitness advantage in marginally suitable environments (Valladares *et al.*, 2014; Larson and Funk, 2016). In the context of long-lived tree species, which are likely to evolve at a slower pace than the more rapid rate of climate change (Savolainen *et al.*, 2007), the degree of plasticity in trait expression is critical for future persistence (Rehfeldt *et al.*, 2002; Aitken *et al.*, 2008; Oddou-Muratorio and Davi, 2014).

Pinus monophylla is a foundational tree species that occupies a large geographical area in the western USA, spanning broad climatic gradients across elevation and latitude (Ziffer-Berger *et al.*, 2014). Recently, a regional drought between 2013 and 2015 in the Great Basin caused extensive dieback of this species (Flake and Weisberg, 2019). Recruitment of *P. monophylla* seedlings is expected to impose a bottleneck period for woodland stand recovery due to short-lived seeds that are dependent on favourable growing seasons and lack of available nurse shrubs for critical microhabitat (Miller *et al.*, 2019; Urza *et al.*, 2019). While previous research has shown that adult trait variation (e.g. seed size and needle morphology) corresponds to regional-scale climate gradients (Vasey *et al.*, 2022), we currently lack basic knowledge on seedling phenotypic trait variation for *P. monophylla*, and on whether seedling responses to drought differ among seed sources.

To this aim, we conducted a common garden glasshouse experiment for *P. monophylla* to explore how seedling growth strategies respond to water availability. Seedlings were propagated from seeds collected from 23 elevationally distributed sites within nine distinct mountain ranges that encompassed the large gradients of temperature and seasonality of precipitation across *P. monophylla*’s range. We applied four watering treatments, representing a drought-stress gradient, to the seedlings and measured above- and below-ground growth-related traits and biomass allocation ratios. We quantified trait plasticity across watering treatments by comparing seedling offspring from each maternal tree. We asked the following questions: (i) How do functional traits and biomass allocation ratios of seedlings from different seed source environments respond to a gradient of water availability? (ii) How is phenotypic trait plasticity, as indicated by response to water availability, related to seed source environment?

We hypothesized that the trait values and plasticity of growth-related seedling traits would show patterns consistent with local adaptation, given clinal variation among seed source environments. For our first question, we predicted that seedlings from drier sites (high water deficit) would respond to moisture-limited treatments by growing longer and heavier roots and producing less above-ground biomass than seedlings from more mesic sites, a strategy that can reduce water stress. Conversely, we predicted that seedlings from wetter sites (both summer-wet and winter-wet) would invest more in above-ground growth than seedlings from drier sites when grown in wetter treatments. Seedlings from wetter sites, adapted to longer periods of growing season water availability, were predicted to produce greater leaf area, which would allow them to assimilate more carbon. For our second question on trait plasticity, we predicted that seedlings from hotter and drier environments and from sites that experience more temporal variation in environmental conditions, such as those that experience episodic pulses of summer precipitation, would exhibit more plasticity. We expected these seedlings to have greater capacity to shift growth strategies in response to water availability, from rapid growth rates in the well-watered treatments to slower, more conservative growth strategies in the moisture-limited treatments. An understanding of trait variation and plasticity in seedling responses to water availability provides information on probable woodland recovery following drought-induced overstorey mortality events across the range of *P. monophylla*.

MATERIALS AND METHODS

Study species, seed collection and seed stratification

Pinus monophylla, or singleleaf pinyon pine, is a widespread dryland tree species of the Great Basin, extending into uppermontane environments of the adjoining Mojave Desert. *Pinus monophylla* often co-occurs with *Juniperus* tree species, and pinyon–juniper woodlands are one of the most important vegetation types within the western USA, occupying >100 million acres (Romme *et al.*, 2009). *Pinus monophylla* generally ranges in elevation between 1000 and 2800 m, with low-elevation sites neighbouring shrubland and grassland communities, and high-elevation sites intermixing with mesic shrubland and subalpine

forest communities (Meeuwig *et al.*, 1990). As one of the most drought-tolerant tree species globally, it grows in areas that receive 200–460 mm of precipitation annually (Meeuwig *et al.*, 1990). It is considered a foundation species, providing habitat and food for many animals, critical ecosystem structure, and primary productivity (Lanner, 1981).

To investigate trait variation across the largest possible range of environmental conditions for *P. monophylla*, we stratified our sampling locations using PRISM 30-year climate normals (Daly *et al.*, 1994) of three mean annual precipitation \times three monsoonal categories. We then selected nine mountain ranges that represented each combination of these categories and were distributed relatively evenly in geographical space (Fig. 1; Vasey *et al.*, 2022) (Supplementary Data Table S1). At each mountain range, we aimed to establish three sampling sites along an elevational gradient, separated by at least 200 m (low, middle and high). The low-elevation site for each mountain range was selected based on first detection of cone-bearing *P. monophylla* trees, while the high-elevation site represented

the highest elevation of cone-bearing trees accessible from a road. The middle-elevation site was located at the elevational midpoint between the other two sites. In three of the mountain ranges, sampling was limited to one or two sites due to narrow elevation ranges or a lack of available cones. The sampling site elevations varied from 1515 to 2611 m, with a total of 23 sites (Fig. 1; Table S1).

At each site, we selected six maternal trees representative of the stand (based on tree size and cone production), and for each tree we harvested at least 30 mature cones in autumn 2019. After the cones opened, we removed the seeds and tested for viability using the acoustic method, following Chambers (2001). We then weighed 20 seeds (fresh weight) per tree to determine mean maternal seed mass. Up to 70 seeds per maternal tree were put into cold, moist stratification for 65 d. Seeds were flushed with running water for 36 h to imbibe the seeds (Kildisheva *et al.*, 2012) and rinse away pathogens (James and Genz, 1981), then placed in plastic bags with wetted vermiculite and stratified in a refrigerator with an average temperature

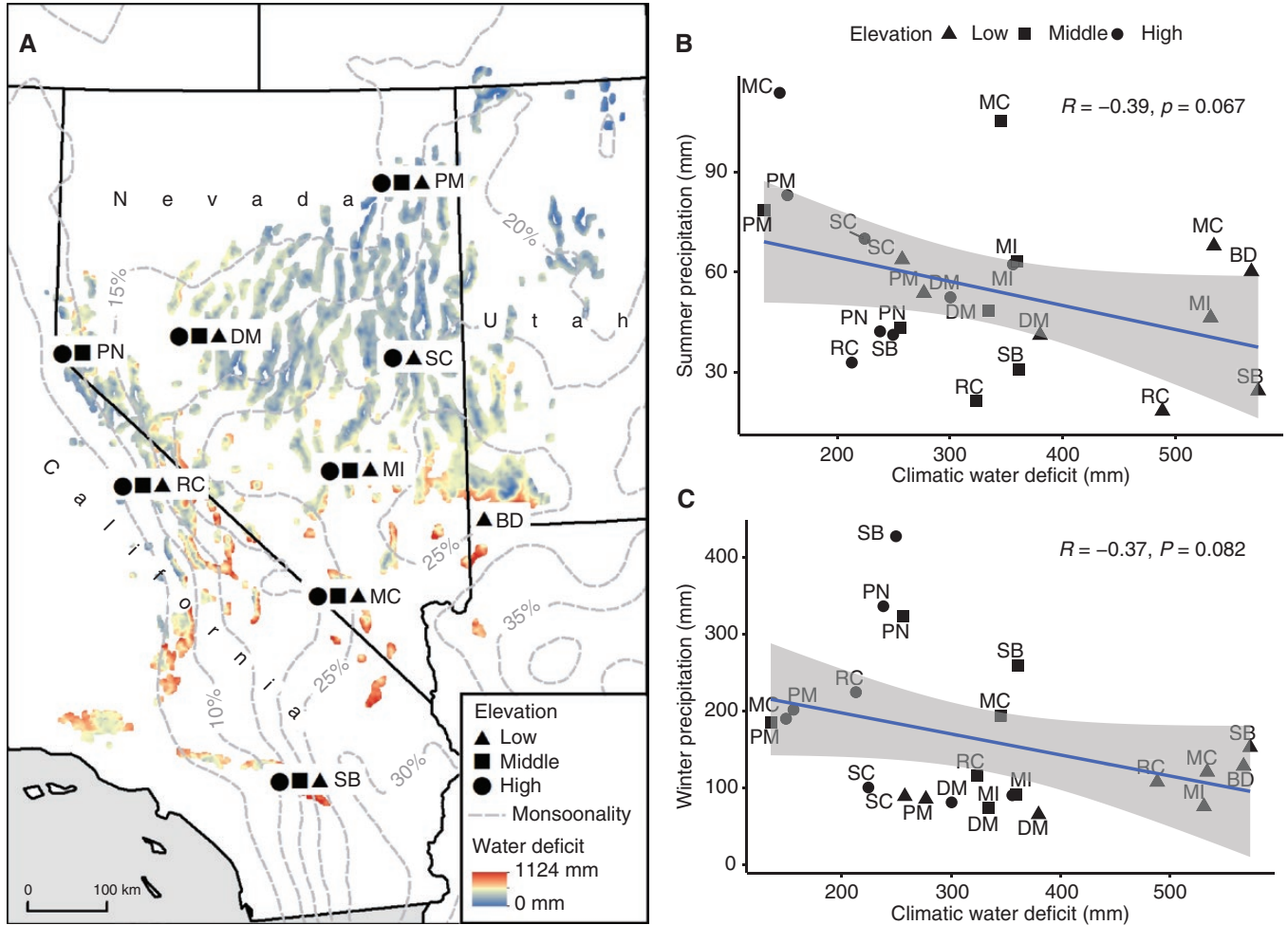


FIG. 1. (A) Seed collection sites. Seeds were collected from 23 elevationally distributed sites (symbols) in nine mountain ranges (two-letter codes: Pine Nut Mountains 'PN', Desatoya Mountains 'DM', Pequot Mountains 'PM', Schell Creek Mountains 'SC', Beaver Dam Mountains 'BD', Mount Irish 'MI', Mount Charleston 'MC', San Bernardino Mountains 'SB' and Rock Creek 'RC'). *Pinus monophylla* species distribution shown in shaded areas, coloured according to climatic water deficit (red = higher water deficit; blue = lower water deficit), overlaid with contour lines representing monsoinality (i.e. the percentage of annual precipitation received in July, August and September). (B) Relationship between 30-year normals of summer precipitation and cumulative climatic water deficit for the seed collection sites. (C) Relationship between 30-year normals of winter precipitation and cumulative climatic water deficit for the seed collection sites.

of 5 °C. Seeds were checked weekly, and when 10 % of the seeds had grown radicles, planting was initiated in the common garden glasshouse experiment in February 2020.

Common garden experimental design and glasshouse methods

Our common garden experiment took place in a climate-controlled glasshouse in Reno, NV, USA. We used a randomized block experimental design, with four watering treatments and three replicate blocks of each treatment for a total of 12 treatment blocks. Each block contained two or three replicates of stratified seeds from each of the 137 maternal trees ($N = 3740$). Each block had 300–320 planting cells consisting of 2.7-inch-diameter \times 14-inch-tall heavy deepot cells (D60H; Stuewe & Sons, Inc.) filled with a locally obtained potting mix that was selected as being similar to native pinyon–juniper woodland soils (Triple mix; Reno Rock, Reno, NV, USA). Two stratified seeds from a randomly assigned maternal tree and seed collection site were planted into each cell to a depth of 3 cm (Chambers, 2001).

Seedlings were grown in four watering treatments that represented a gradient of soil water availability (Supplementary Data Fig. S1). To assign the watering treatments, we used a gravimetric method to track soil water content (Dumroese *et al.*, 2015). We first selected five filled containers per treatment block that were of comparable weight at saturation (within 10 g). We then dried the soil from one of the five containers at 110 °C to obtain the oven-dry soil weight, which we used to convert the remaining deepot cells' weight to gravimetric water content (GWC). We developed a soil water release curve for six fine soil (0.065 mm) subsamples across a range of known GWCs, allowing us to determine field capacity (maximum saturation) at 40 % GWC and permanent wilting point (-1.5 MPa) at 10 % GWC (WP4C water potential meter). All seedlings were given the wettest treatment for 1.5 months until most of the seedlings had emerged, after which we used a gradual dry-down process to achieve target GWC ranges for the watering treatments: 'Wettest' (34–40 % GWC), 'Wet' (22–28 % GWC), 'Dry' (10–16 % GWC) and 'Driest' (no addition of water following the initial seedling emergence period). Selected deepot cells were weighed daily, and irrigation was initiated for the different watering treatments when they reached the low end of the assigned GWC range (Fig. S1). Seedlings were irrigated with inverted rotor spray mini sprinklers using a fine mist. For deepot cells that had two seedlings emerge, we removed one of the seedlings to avoid competition. Trays were rotated within treatment blocks before each watering treatment to account for minor variability within each block. Temperatures in the glasshouse were maintained between 12 and 27 °C, within the diurnal range of summer temperatures experienced by naturally occurring *P. monophylla* seedlings across an elevational gradient (Urza *et al.*, 2019). The experiment was concluded at the end of August 2020, following 4.5 months of treatment and 6 months of total growth.

Measurements of morphological and performance traits

Biweekly checks for status (not emerged, emerged and dead) began 3 weeks following planting at the first sign of emergence,

and continued for the rest of the experiment. In total, 3320 seedlings emerged and survived, 91 emerged and died, and 329 never emerged.

Thirteen phenotypic traits were measured at the end of the experiment. Above-ground traits included height (mm), canopy area (mm²), stem diameter (mm), above-ground dried biomass (g) and number of branches. Below-ground traits included dried root biomass (g), root length (cm), specific root length (m g⁻¹), root volume (cm³) and root surface area (cm²). We calculated three biomass allocation ratios (Poorter *et al.*, 2012) from the above- and below-ground dry weights, including needle mass fraction (NMF, ratio of leaf biomass to total biomass of the seedling), stem mass fraction (SMF, ratio of stem biomass to total biomass of the seedling) and root mass fraction (RMF, ratio of root biomass to total biomass of seedling).

Height, canopy area and number of branches were measured at the end of the experiment for all surviving seedlings ($N = 3320$). Height was measured using a transparent metric ruler (1-mm precision) from the soil surface to the top of the plant. Canopy diameter at its widest point (d^a) and perpendicular to its widest point (d^b) was measured using a ruler (1-mm precision), and canopy area was calculated using the equation for an ellipse. Number of branches was counted as the number of lateral shoots growing off the main stem.

Biomass was measured using destructive methods for approximately one-third of the seedlings ($N = 1113$). Following root extraction and cleaning, stem diameter was measured at the level of the soil surface using a digital caliper, and above- and below-ground tissues were separated. Using a dual-light flatbed scanner (Epson Expression 1100XL; Seiko Epson, Suwa, Japan), fresh roots were scanned to obtain high-resolution imagery (1200 dpi) to measure root length, root volume and root surface area (WinRhizo; Regent Instruments, Quebec, Canada). Above- and below-ground tissues were dried at 70 °C for 4 h prior to weighing (100 g \times 0.001 g scale). Needles, twigs and stems were separated and weighed individually to obtain the different above-ground biomass components for biomass allocation ratios. To calculate specific root length, root length was divided by root biomass for each seedling.

Environmental variables

Four environmental variables were derived for each seed collection site from existing spatial data layers: cumulative climatic water deficit (CumlCWD, mm), soil available water storage capacity (AWC, cm), winter precipitation (mm) and summer precipitation (mm) (Supplementary Data Table S1). Correlations among these variables were <0.41 . In the Intermountain West, these environmental variables are strongly associated with water balance and help to define the lower climatic tree limit of pinyon and juniper woodlands (Urza *et al.*, 2020), and they are correlated with *P. monophylla* adult tree traits (Vasey *et al.*, 2022).

CumlCWD was calculated using a Thornthwaite water balance model (Lutz *et al.*, 2010), which incorporates 30-year monthly normals (800-m cell size) of temperature and precipitation, elevation, and soil water holding capacity, using methods described by Dilts *et al.* (2015). Often considered a proxy for aridity, CumlCWD is calculated as the difference

between potential evapotranspiration and actual evapotranspiration (Stephenson, 1990). It represents the evaporative demand not met by available water. In our Great Basin study area, CumlCWD is negatively associated with increasing elevation (i.e. higher elevations are wetter by this metric; Dilts *et al.*, 2015).

AWC was obtained from the POLARIS soil map at 90-m cell size resolution (Chaney *et al.*, 2016). Low AWC is generally associated with coarser, sandy soils with increased permeability. Water in low-AWC soils thus collects in deep soil pools (Noy-Meir, 1973), which is potentially problematic for seedlings with small roots (Redmond and Barger, 2013).

Winter precipitation (sum of precipitation in December, January and February) and summer precipitation (sum of precipitation in June, July and August) were calculated using 30-year normal (1981–2010) gridded PRISM climate data (PRISM Climate Group 2015) with 800-m cell size (Daly *et al.*, 1994). Seasonal precipitation patterns vary across the range of *P. monophylla*: populations in western and northwestern areas receive most of their annual precipitation as winter snow, whereas populations in more eastern and southeastern areas experience both winter and summer precipitation due to weak monsoonal patterns (Adams and Comrie, 1997). This difference in timing affects the seasonality of growth. Populations depending exclusively on winter snowmelt grow from late spring to early summer, whereas populations receiving summer precipitation can have bimodal patterns of growth between late spring and early summer, and then again from late summer to early autumn (Notaro *et al.*, 2010).

Statistical analysis

Modelling of traits. To evaluate how seedling growth-related traits varied in response to watering treatment and seed source environment, we assessed each trait using linear mixed-effects models fit with maximum likelihood using the ‘lme4’ package in R Studio (Bates *et al.*, 2015; R Core Team, 2020). The fixed effects included watering treatment, seed source environmental variables, and the interaction of treatment and seed source environment. To control for differences at the start of the experiment, we also included fixed effects for seedling emergence time (in days) and mean seed mass for each maternal tree. The random effects included block, maternal tree and mountain range. All fixed effects were standardized using *z*-scores. To evaluate the statistical significance of the fixed effects, we calculated analysis of variance tables with *F*-tests and *P*-values using Satterthwaite approximated degrees of freedom using the package ‘lmerTest’ (Kuznetsova *et al.*, 2017). To visualize the marginal effects of seed source environmental variables, or predictions generated by the model when holding the non-focal variables constant and varying the focal variable, we used the ‘ggpredict’ function from the ‘ggeffects’ package in R (Lüdtke, 2018) and plotted results using ggplot2 (Wickham, 2016).

Calculating and modelling phenotypic plasticity. Because the effect of watering treatment was non-linear, we evaluated plasticity as a trait using a relativized metric, the Relative Distance Plasticity Index (RDPI) (Valladares *et al.*, 2006). We computed

trait plasticity for each maternal tree by calculating the relative trait distance between individual seedlings for each pairwise treatment combination. This allowed us to quantify drought-related phenotypic plasticities for each variable as independent traits that varied between different source environments.

To calculate the RDPI values for each maternal tree, we used the ‘Plasticity’ package in R (Ameztegui, 2017). Each phenotypic distance was calculated by taking the absolute value of the difference between phenotypic trait values of seedlings from the same maternal trees in different watering treatments divided by the sum of those phenotypic trait values [e.g. $|Trait_{Wettest} - Trait_{Wet}| / (Trait_{Wettest} + Trait_{Wet})$]. This was calculated for all combinations of treatment pairs, and final RDPI values were calculated as the average of the normalized treatment differences for each maternal tree. This non-parametric index provides a relativized value that ranges between 0 (no plasticity) and 1 (maximal plasticity), providing a consistent basis for comparison across different populations and traits.

From these outputs, we developed linear mixed-effects models with RDPI of each trait as the dependent variable as a function of fixed effects (seed source environmental variables, emergence time and seed mass) and random effects (maternal tree and mountain range). We used the same R packages for the plasticity analysis as were used in the trait analysis.

RESULTS

Above-ground traits, below-ground traits and allocation ratios

After accounting for seed mass and emergence time, seedling traits and allocation ratios remained strongly influenced by the environmental variables characterizing the seed source habitat (Table 1). Seedlings from drier climates (higher cumulative climatic water deficit), coarse or shallow soils (lower soil available water capacity) and greater precipitation in the summer months (higher summer precipitation) tended to produce larger above-ground structures. Overall, above-ground growth was smallest in the dry and driest watering treatments, while the optimal treatment for above-ground growth was the wet treatment (Supplementary Data Fig. S2: 22–28 % GWC) suggesting that more moisture increases above-ground growth only to a threshold level and that the wettest treatment was probably too moist for seedlings of this drought-tolerant species.

Within all treatment groups, seedling heights were greatest from sites with higher climatic water deficit (Table 1, Fig. 2A). Seedlings receiving the two driest treatments responded similarly to water availability regardless of summer precipitation of the source environment, whereas summer precipitation was positively associated with height growth for seedlings receiving the wettest treatments [Fig. 2B; *p* (summer precipitation × treatment [T]) = 0.01]. Similarly, soil available water capacity of the seed source was most strongly negatively related to seedling height in the wettest treatments [Fig. 2C; *p* (soil available water capacity × T) = 0.03]. Seedling canopy area was largest for seed sources associated with higher climatic water deficit and greater summer precipitation for all treatments (Table 1, Fig. 2D, E), and both seed source environmental variables had significant interactions with treatment (Table 1). Stem diameter was greater for seed sources from higher climatic water deficit

TABLE 1. Linear mixed-effect model outputs (F-test and P-value) for seedling above-ground traits, below-ground traits and biomass allocation ratios. For each trait and fixed effect (watering treatment, environmental conditions in seed source locations, emergence time, seed mass, and the interaction of treatment and seed source environment) the F-statistic is reported in the top row, with P-values listed directly below. P-values <0.05 are in bold type.

Trait	Fixed effects predictions											
Interaction term												
	F-test/ <i>P</i> -value	Watering treatment (TRT)	Climatic water deficit	Summer precipitation	Winter precipitation	Available water storage capacity	Emergence time	Seed weight	Climatic water deficit × TRT	Summer precipitation × TRT	Winter precipitation × TRT	Available water storage × TRT
Above-ground												
Height (mm)	<i>F</i>	49.6	4.2	2.2	1.2	1.6	14.8	31.3	1.6	4.2	2.2	3.1
	<i>P</i>	0.001	0.05	0.16	0.28	0.21	0.001	0.001	0.19	0.01	0.09	0.03
Canopy area (mm ²)	<i>F</i>	1.5	14.4	10.6	4.8	0.2	28.6	28.4	4.3	4.6	2.0	0.3
	<i>P</i>	0.28	0.001	0.001	0.03	0.66	0.001	0.001	0.001	0.001	0.12	0.85
Stem diameter (mm)	<i>F</i>	267.9	30.8	9.8	0.8	1.8	28.9	99.6	0.7	0.1	0.1	0.9
	<i>P</i>	0.001	0.001	0.04	0.41	0.20	0.001	0.001	0.54	0.95	0.93	0.43
Above-ground biomass (g)	<i>F</i>	316.7	8.9	5.7	0.1	2.5	43.3	65.3	1.9	1.0	1.1	2.1
	<i>P</i>	0.001	0.01	0.06	0.75	0.14	0.001	0.001	0.12	0.38	0.34	0.09
Number of branches	<i>F</i>	4.3	1.3	13.6	1.9	2.2	16.0	8.3	4.0	9.4	1.2	1.9
	<i>P</i>	0.03	0.26	0.001	0.19	0.15	0.001	0.001	0.01	0.001	0.31	0.13
Below-ground												
Root biomass (log: g)	<i>F</i>	269.4	8.2	1.8	2.6	0.8	35.3	120.2	1.37	0.3	0.6	0.1
	<i>P</i>	0.001	0.001	0.18	0.11	0.39	0.001	0.001	0.25	0.82	0.62	0.97
Root length (cm)	<i>F</i>	231.3	0.1	0.1	1.7	0.00	37.4	41.9	1.1	2.2	1.8	0.1
	<i>P</i>	0.001	0.77	0.74	0.19	1.00	0.001	0.001	0.33	0.09	0.14	0.97
Specific root length (m g ⁻¹)	<i>F</i>	26.0	11.9	3.4	2.0	0.2	0.3	22.3	0.6	2.0	1.1	0.7
	<i>P</i>	0.001	0.001	0.12	0.19	0.70	0.57	0.001	0.60	0.11	0.35	0.58
Root volume (log: cm ³)	<i>F</i>	446.6	2.7	1.8	2.7	1.1	23.5	102.7	1.5	0.7	1.5	0.1
	<i>P</i>	0.001	0.10	0.26	0.11	0.30	0.001	0.001	0.21	0.57	0.21	0.98
Root surface area (cm ²)	<i>F</i>	212.6	0.0	0.0	3.3	0.0	28.0	37.8	0.9	1.7	1.7	0.1
	<i>P</i>	0.001	0.90	0.91	0.07	0.86	0.001	0.001	0.43	0.16	0.16	0.98
Allocation ratios												
Needle mass fraction	<i>F</i>	48.8	5.7	1.4	1.0	5.6	2.3	4.5	0.4	2.4	0.5	2.0
	<i>P</i>	0.001	0.02	0.27	0.34	0.03	0.13	0.04	0.77	0.07	0.66	0.11
Stem mass fraction (log)	<i>F</i>	160.2	0.1	0.5	0.8	0.3	16.2	0.0	0.2	0.5	1.5	0.6
	<i>P</i>	0.001	0.82	0.48	0.38	0.57	0.001	0.86	0.93	0.65	0.21	0.61
Root mass fraction	<i>F</i>	77.5	1.4	2.0	0.9	0.1	0.3	5.4	1.1	1.3	0.3	0.9
	<i>P</i>	0.001	0.25	0.19	0.35	0.80	0.61	0.02	0.37	0.29	0.84	0.45

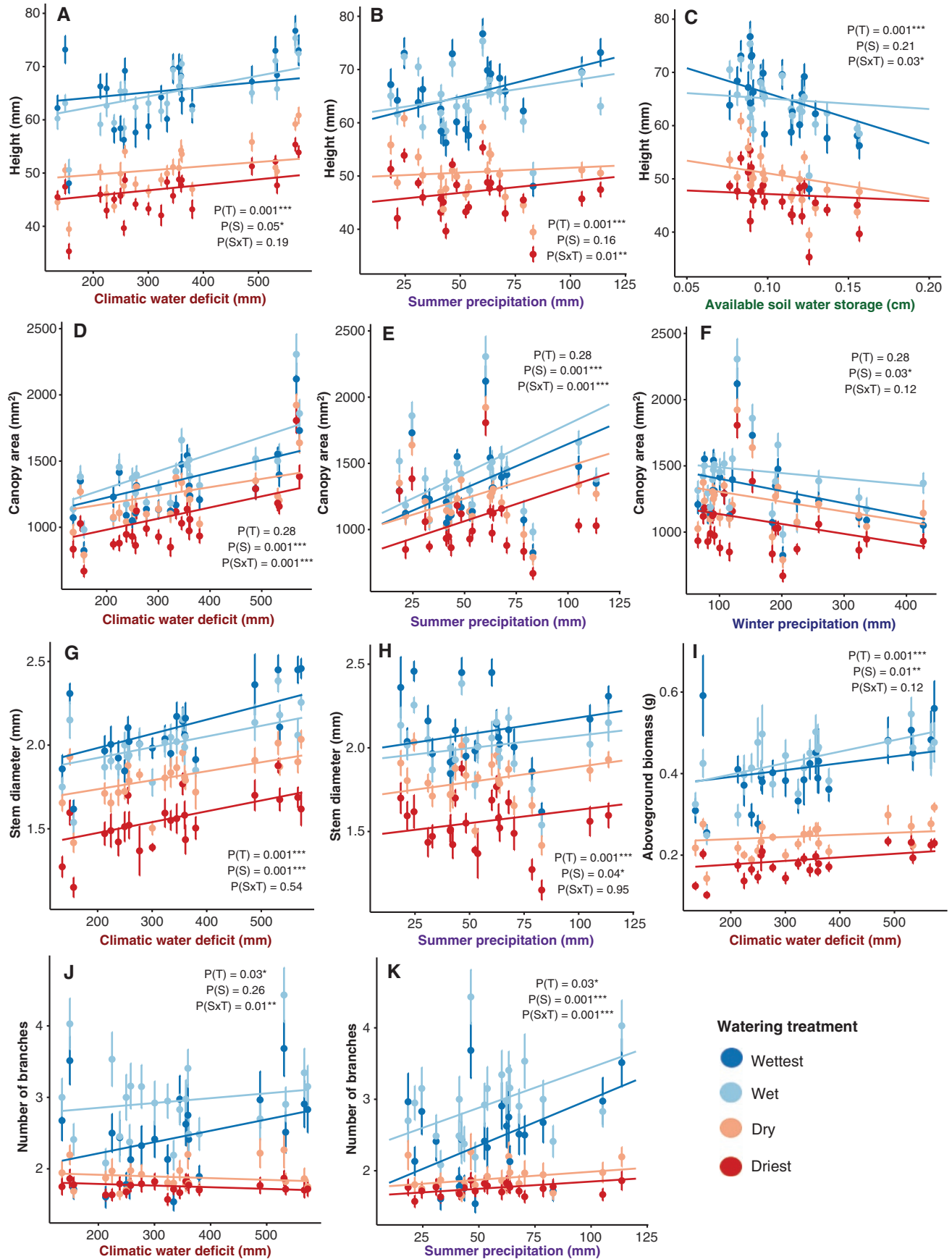


FIG. 2. Partial effects plots for models of above-ground seedling trait values. Points represent mean trait values and standard errors for each seed source as a function of watering treatment (T) and seed source environmental conditions (S). Lines represent the partial effects of the environmental variable and its interaction with watering treatment. The four colours represent the different watering treatments: wettest = dark blue, wet = light blue, dry = peach and driest = red. Only traits with significant ($P < 0.05$) effects of seed source environment (S) or seed source environment \times treatment (S \times T) interactions are shown.

and summer precipitation under all treatments (Table 1, Fig. 2G, H). Seedling total above-ground biomass was greater with seed sources from higher climatic water deficit (Table 1, Fig. 2I). Seedlings from drier sources (higher climatic water deficit) grew more branches than seedlings from wetter sources in the two wet treatments, but seedlings from all seed sources grew similar numbers of branches in the two dry treatments (Table 1, Fig. 2J). The dependence of branches on seed source was even stronger for summer precipitation, but, in this case, seedlings with wetter summers (higher summer precipitation) grew more branches in the wet treatments than seedlings with drier summers (Table 1, Fig. 2K).

Root traits had fewer significant relationships with seed source environmental variables overall, and source environment and watering treatment had no significant interactions. However, all root traits were significantly affected by watering treatment (Table 1), decreasing in growth under the drier watering treatments (Supplementary Data Fig. S3). Seedling root biomass was greater for seed sources with higher climatic water deficit in all treatments (Table 1, Fig. 3A). Conversely, specific root length decreased for seed sources with higher climatic water deficit (Table 1, Fig. 3B).

Although all biomass allocation ratios were significantly affected by watering treatment, only needle mass fraction had significant seed source environmental effects (Table 1, Fig. 4). Needle mass fraction had a weak positive relationship with climatic water deficit and soil available water capacity of the seed source environment (Table 1, Fig. 4A, B). Overall, the drier watering treatments produced seedlings with lower needle mass and with higher stem mass fraction and root mass fraction (Supplementary Data Fig. S4). None of the biomass allocation ratios had significant interactions with seed source environment and watering treatment.

Plasticity

Plasticity in certain traits varied depending on the seed source environment and was most strongly related to the magnitude of summer precipitation (Supplementary Data Figs S5–S7). Above-ground biomass plasticity ranged from 0 to 0.92 and was greatest for seed sources with greater climatic water deficit, summer precipitation and winter precipitation, with the strongest effects of summer precipitation (Table 2, Fig. 5A–C). Plasticity in number of branches (0–1) was greatest for seed sources with greater summer precipitation (Table 2, Fig. 5D). Plasticity in biomass allocation ratios also did not vary with seed source environment (Table 2; Fig. S7).

Interestingly, although root mean traits showed little variation with seed source environment, root trait plasticity showed consistent patterns of variation (Supplementary Data Fig. S6). Root biomass plasticity (0–0.86) was greatest for seed sources with higher climatic water deficit and greater summer precipitation (Table 1, Fig. 5E, F). Root length plasticity (0–0.92) was greatest for seed sources with greater summer precipitation (Table 2, Fig. 5G). Lastly, root surface area plasticity (0–0.86) was greatest for seed sources with greater summer precipitation (Table 2, Fig. 5H). Trait plasticity in response to watering treatments did not vary significantly with soil available water capacity of the seed source (Table 2).

DISCUSSION

Seedling trait values vary along seed source environmental gradients

Our findings showed that seedling trait values varied with watering treatment, and that these responses differed depending on the environmental conditions associated with the seed sources. Contrary to our predictions, seedlings sourced from drier climates (higher cumulative climatic water deficit) or from sites with greater summer precipitation grew more above ground in all treatments compared to seedlings sourced from wetter sites. In general, the influence of water deficit on above-ground seedling traits was of greater magnitude than the influence of summer precipitation. Root traits followed the expected trend, with seedlings from drier climates producing higher root biomass and lower specific root length than seedlings from wetter climates. Notably, seedling growth was strongly related to climatic water deficit even after accounting for the effect of seed mass, which for *P. monophylla* has been shown to be greater in more arid sites (Vasey et al., 2022). Overall, clinal variation in traits that correspond to environmental gradients is consistent with local adaptation, supporting our hypothesis.

Our results suggest that seedlings from areas with greater water deficits, where good water years are scarce, have developed adaptations that enable greater above- and below-ground growth when water is available. Seedlings from drier environments grew taller, with larger canopy area, thicker stems and greater root biomass, compared to seedlings from more mesic environments. Contrary to our prediction that seedlings from the drier seed sources would only establish larger root systems, we found that overall growth of these seedlings was greater both above- and below-ground than seedlings from wetter environments. This strategy of rapid growth can increase carbon gain at the start of establishment and help ensure resource reserves for later use (Li et al., 2020), while also producing more extensive root systems to minimize drought stress in the long run. Allocating more resources to above-ground biomass, however, can increase the risk of mortality (e.g. hydraulic failure), particularly when resources become limited (Augustine and Reinhardt, 2019; Shen et al., 2019).

It is adaptive for seedlings under drought stress to increase root biomass via root diameter (i.e. thicker roots), a strategy that can increase survival in dry environments, defend against pests and pathogens, and maintain transport capacity and stability (Poorter and Ryser, 2015; Larson and Funk, 2016; Shen et al., 2019). High specific root length, or having long, thin roots, is less expensive to produce and allows greater resource acquisition (i.e. greater absorptive length per unit biomass), but comes at the cost of being less stress-tolerant (Withington et al., 2006; Larson and Funk, 2016). We found that seedlings sourced from drier locations within the range of *P. monophylla* consistently had the greatest root biomass and lower specific root length, potentially allowing seedlings to withstand greater environmental stress. In contrast, seedlings sourced from wetter locations generally had thinner and longer roots that are well suited for resource acquisition during the growing season when moisture is available. However, the relationship between root biomass and seed source environment was weaker in the driest watering treatments, suggesting that the capacity for seedlings

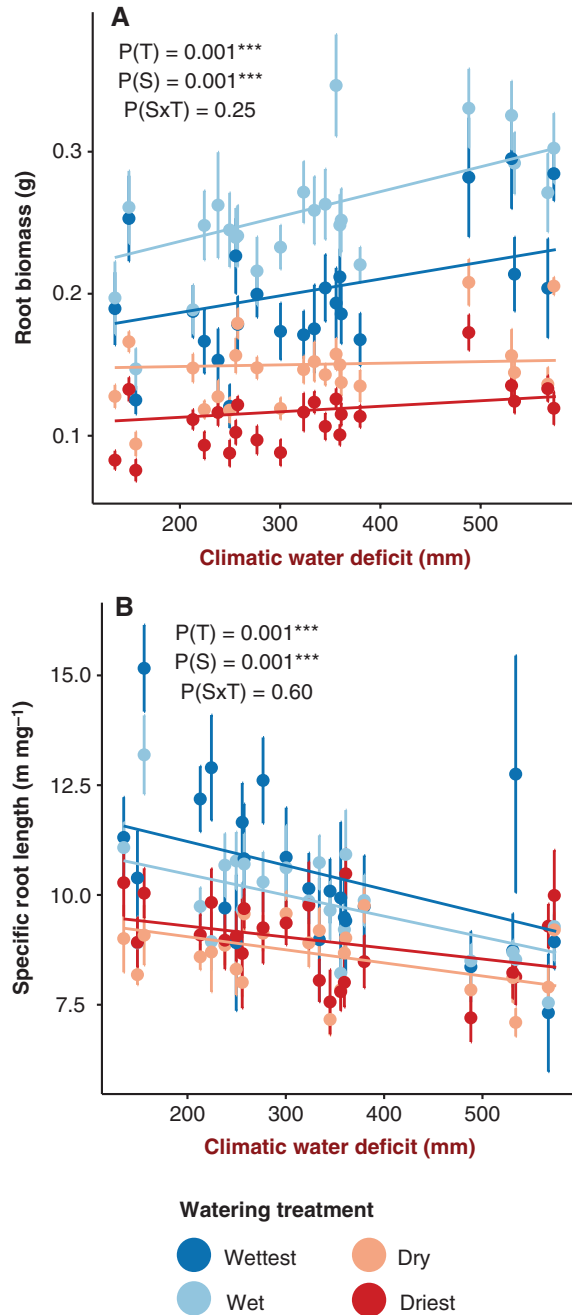


FIG. 3. Partial effects plots for models of below-ground seedling trait values. Points represent mean trait values and standard errors for each seed source as a function of watering treatment (T) and seed source environmental conditions (S). Lines represent the partial effects of the environmental variable and its interaction with watering treatment. The four colours represent the different watering treatments: wettest = dark blue, wet = light blue, dry = peach and driest = red. Only traits with significant ($P < 0.05$) effects of seed source environment (S) or seed source environment \times treatment ($S \times T$) interactions are shown.

to express phenotypic variation may be reduced under extreme stress (Diethelm *et al.*, 2022).

Consistent with prior studies (Jose *et al.*, 2003; Padilla *et al.*, 2007; Poorter *et al.*, 2012), our study found that under watering treatments that simulate extreme drought conditions, seedlings

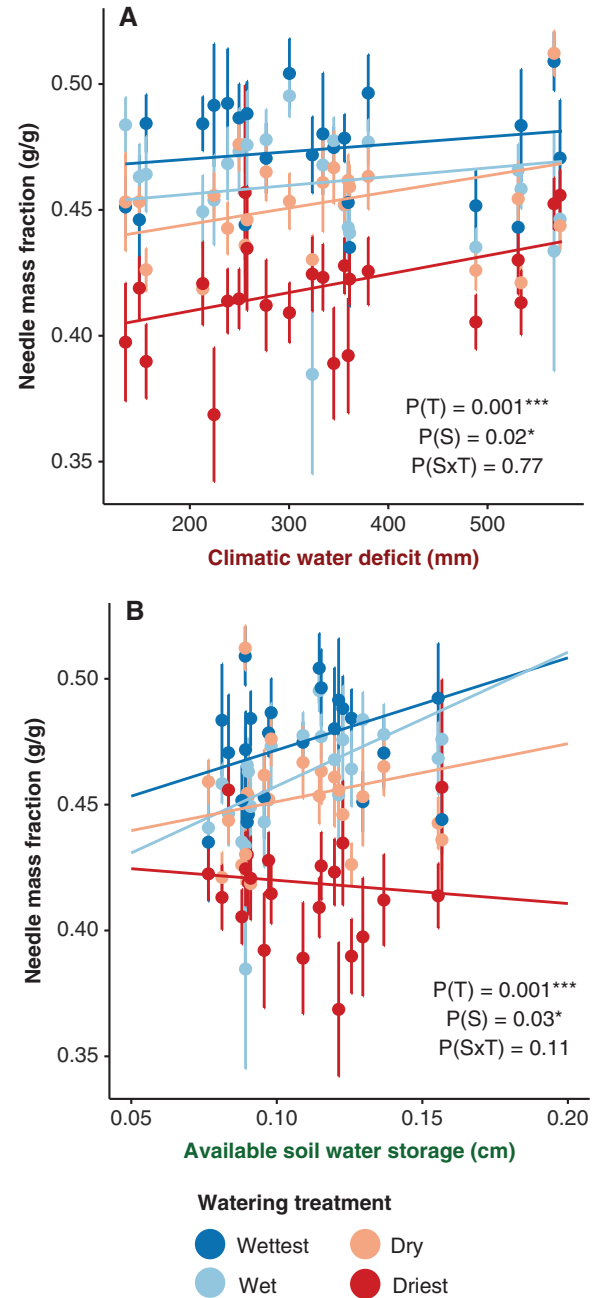


FIG. 4. Partial effects plots for models of seedling biomass allocation ratios. Points represent mean trait values and standard errors for each seed source as a function of watering treatment (T) and seed source environmental conditions (S). Lines represent the partial effects of the environmental variable and its interaction with watering treatment. The four colours represent the different watering treatments: wettest = dark blue, wet = light blue, dry = peach and driest = red. Only traits with significant ($P < 0.05$) effects of seed source environment (S) or seed source environment \times treatment ($S \times T$) interactions are shown.

increased their biomass allocations to stems and roots while reducing their needle allocation regardless of seed source environment. Because shoot and root biomass showed similar relationships across seed source environments, biomass allocation ratios responded primarily to watering treatment, and the

TABLE 2. Linear mixed-effects model outputs for RDPI values as the dependent variables. For each trait the F-statistic is shown in the top row and the associated P-value directly below. P-values <0.05 are in bold type. Trait plasticity in six models had significant relationships with seed source environment.

Trait	Fixed effects predictions					
	F-test/P-value	Climatic water deficit	Summer precipitation	Winter precipitation	Available soil water storage	Emergence time
Above-ground						
Height plasticity	F	0.0	2.5	0.0	0.9	1.2
	P	0.99	0.12	0.96	0.36	0.58
Canopy area plasticity	F	2.6	0.4	1.5	0.5	0.6
	P	0.11	0.51	0.23	0.49	0.45
Stem diameter plasticity	F	0.0	0.1	0.0	0.1	1.9
	P	0.97	0.76	0.93	0.73	0.17
Total above-ground biomass plasticity	F	3.9	16.5	4.2	0.1	0.8
	P	0.05	<0.001	0.04	0.74	0.36
Number of branches plasticity	F	0.2	9.9	1.0	1.6	0.2
	P	0.70	<0.001	0.33	0.21	0.70
Below-ground						
Root biomass plasticity	F	7.1	7.1	3.2	2.3	2.9
	P	0.01	0.01	0.08	0.13	0.09
Root length plasticity	F	2.4	7.1	4.5	1.5	0.0
	P	0.13	0.03	0.06	0.24	0.99
Specific root length plasticity	F	0.3	1.1	1.1	0.6	1.8
	P	0.6	0.3	0.3	0.4	0.2
Root volume plasticity	F	0.00	2.96	2.31	0.45	1.40
	P	1.0	0.1	0.2	0.5	0.2
Root surface area plasticity	F	0.9	5.5	4.4	1.5	0.4
	P	0.35	0.04	0.06	0.24	0.54
Allocation ratios						
Needle mass fraction plasticity	F	0.6	0.1	0.1	0.1	0.9
	P	0.43	0.71	0.81	0.72	0.33
Stem mass fraction plasticity	F	2.3	3.1	0.0	0.1	0.1
	P	0.13	0.08	0.92	0.75	0.77
Root mass fraction plasticity	F	0.4	0.3	2.5	0.5	0.8
	P	0.53	0.57	0.12	0.49	0.38

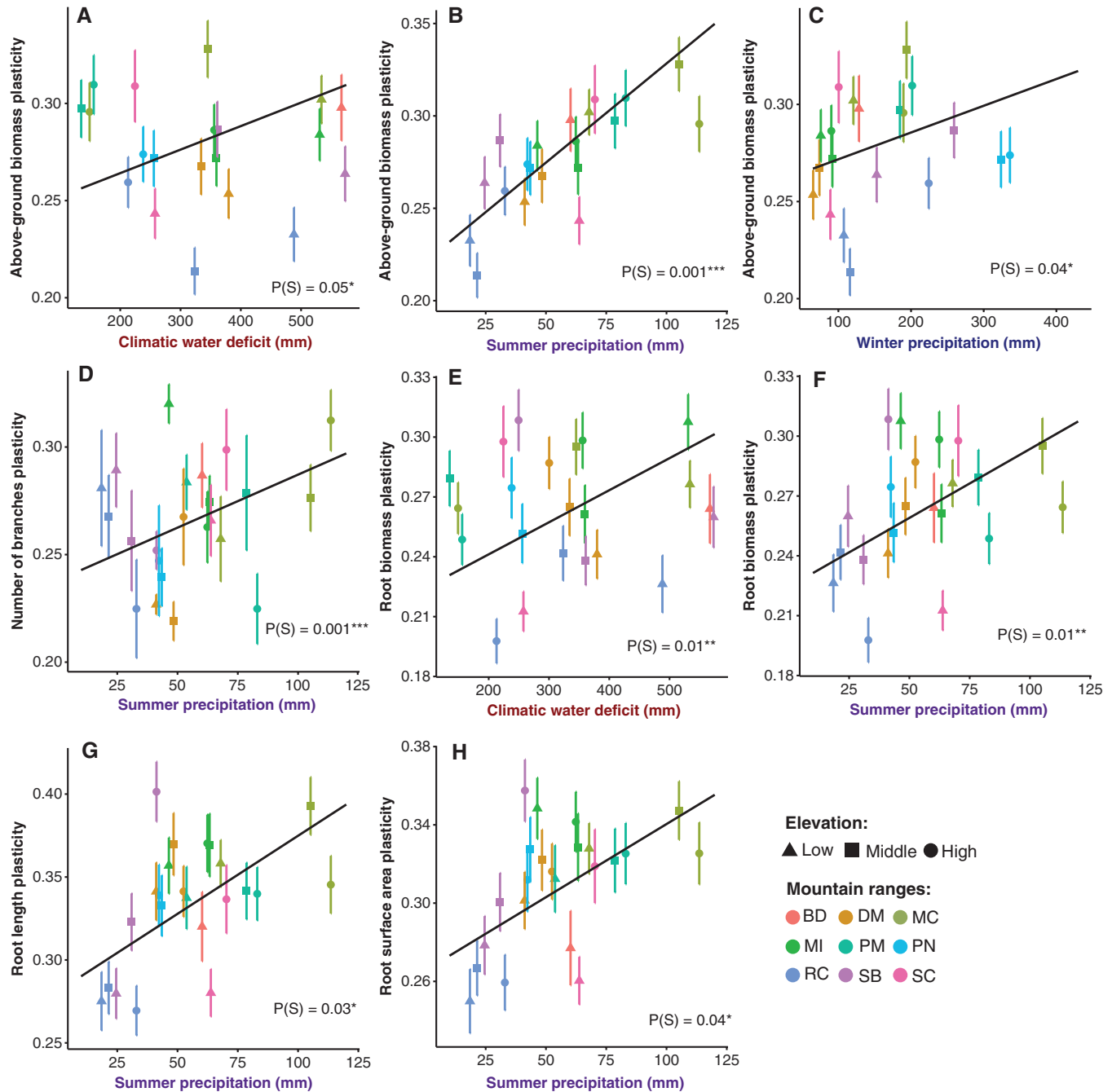


FIG. 5. Partial effects plots for models of trait plasticity in response to watering treatments (calculated as RDPI). Points represent mean plasticity values and standard errors for each seed source as a function of seed source environmental conditions (S). Lines represent the partial effect of the environmental variable. Colours represent mountain ranges, and symbols represent elevation categories for each site. Only traits with significant ($P < 0.05$) effects of seed source environment (S) are shown.

proportion of biomass allocated towards shoots vs. roots did not vary as a function of seed source environment. Although we had predicted that seedlings from drier climates would invest more in below-ground biomass than seedlings from wetter climates, low variability in allocation ratios across seed sources was probably due to plant requirements to maintain balanced performance among leaves, stems and roots (Poorter *et al.*, 2012).

Trait plasticity is related to seed source summer precipitation

In general, the results support our prediction that populations which experience more episodic summer precipitation are more plastic in their trait expressions (Pratt and Mooney, 2013). A trade-off exists between optimizing growth during favourable conditions vs. growing more slowly to avoid subsequent mortality when conditions become more stressful later (Grime, 2006; Martinez-Villata *et al.*, 2010). Populations

from across the range of *P. monophylla* face relatively high growing season water deficits; however, some of these populations (i.e. ‘summer-wet’ populations) also experience periodic summer precipitation from monsoonal weather events which, if supplying sufficient moisture, can signal species in arid and semi-arid climates to initiate growth (Noy-Meir, 1973; Gillespie and Loik, 2004; Walck *et al.*, 2011). *Pinus monophylla* is characterized as an isohydric species that loses its ability to assimilate carbon below a leaf water potential of -2 MPa (Nowak *et al.*, 1999). It is therefore important for this species to utilize infrequent periods of increased soil moisture for carbon gain before closing stomata to prevent water loss during extended dry periods (West, 2006). Thus, our finding that seedlings from summer-wet environments exhibited greater growth plasticity in response to watering is consistent with our hypothesis of local adaptation.

Having greater capacity to quickly respond to changing environmental conditions (i.e. higher trait plasticity) can confer a fitness advantage, particularly in environments that are more resource-limited (Valladares *et al.*, 2014; Larson and Funk, 2016). With less developed root systems, tree seedlings in semi-arid environments mainly have access to the shallow surface soil layer, which is subject to more variable soil moisture conditions (Matías *et al.*, 2012). Although nutrients essential for plant growth are concentrated within this soil layer, soil drying due to drought reduces diffusivity and mass flow of nutrients (Ryel *et al.*, 2008), and precipitation-induced nutrient pulses are only available to roots in contact with the surface layers (Hodge, 2004; Ryel *et al.*, 2008). Thus, greater root plasticity at the recruitment stage could favour uptake of essential nutrients during periods of increased soil moisture (Vizcaíno-Palomar *et al.*, 2020). Although we could not evaluate trait-associated differences in seedling survival because few seedlings died during our study, we found that root biomass plasticity, root length plasticity and root surface area plasticity were highest for seedlings from summer-wet environments, potentially allowing a faster growth response to infrequent resource pulses (Ushio *et al.*, 2015; Vizcaíno-Palomar *et al.*, 2020).

CONCLUSIONS

Tree species worldwide are facing increasingly severe and frequent drought events associated with climate change (Allen *et al.*, 2010). Our study provides insights into how drought stress may affect seedling recruitment in different environments throughout the range of a widespread tree species. Variation in trait expression corresponded to seed source environmental conditions and was structured along regional gradients of precipitation seasonality, as well as more local, elevational gradients of water availability.

Local adaptation in seedling traits – suggested by environmentally structured clinal variation in both trait values and trait plasticity – is expected to influence the ability of *P. monophylla* to adapt *in situ* and establish in future hotter and drier environments. Populations from the drier environments within its range are predicted to be disproportionately affected by increased future drought (Anderegg *et al.*, 2019). Our findings suggest that seedlings from these populations have adaptations that may allow them to survive more stressful environmental conditions,

including efficient growth that increases carbon assimilation, resource storage pools and structural support.

We found that summer precipitation was strongly related to seedling trait responses, suggesting that shifts in precipitation seasonality, and not only changes in total annual water availability, may affect *P. monophylla* populations in future climates. Greater trait plasticity of summer-wet seedlings indicates that plastic growth strategies may have evolved from exposure to infrequent periods of increased soil moisture. Phenotypic plasticity involves non-genetic changes in traits, yet plasticity itself is heritable and can vary among populations and species (Scheiner, 1993). While phenotypic plasticity is generally thought to be adaptive for plant populations experiencing rapid environmental change (Nicotra *et al.*, 2010), it has also been suggested that adaptation to changing climate may be inhibited for phenotypically plastic traits (Westerband *et al.*, 2021). Our results suggest that in the short term, recruitment in trailing-edge (e.g. low-elevation) populations from the summer-dry portions of the range of *P. monophylla* may be negatively impacted by the limited capacity of seedlings to respond plastically to increasing aridity. Because slow-growing, dryland tree species such as *P. monophylla* can spend decades in the juvenile stage, the capacity of seedlings for plasticity in response to environmental stress will be particularly important under ongoing climatic change.

SUPPLEMENTARY DATA

Supplementary data are available online at <https://academic.oup.com/aob> and consist of the following.

Table S1: Site information for each seed source. Fig. S1: Graph of watering treatments over the course of the experiment using gravimetric methods. Fig. S2: Partial effects plots for all above-ground traits. Fig. S3: Partial effects plots for all below-ground traits. Fig. S4: Partial effects plots for all biomass allocation ratios. Fig. S5: Partial effects plots for all above-ground trait plasticities. Fig. S6: Partial effects plots for all below-ground trait plasticities. Fig. S7: Partial effects plots for all biomass allocation ratio trait plasticities.

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